Scientific and Technical Information Center

SEARCH REQUEST FORM

August Committee	SEARCH REQUES	/ /
Requester's Full Name:	NC ZARA Exam	niner # : 77518 Date: 11/05
Art Unit: 16.35 Phone	Number: 2- 07-65	Serial Number: 09/735/363
I - antion (Did-/Doom#), In 28	(Mailbox:#) > C / X Results	Format Preferred (circle): PAPER DISK
********	********	************
To ensure an efficient and quality search,	please attach a copy of the cover sheet	t, claims, and abstract or fill out the following:
• •	1-30	
Title of Invention:	peuling 1	seful systeti deza
Inventors (please provide full names):		to.
·	·	
Earliest Priority Date:	12/00	
Search Topic: Please provide a detailed statement of the se elected species or structures, keywords, syno Define any terms that may have a special m	mums, acronums, and registry numbers	as possible the subject matter to be searched. Include the s, and combine with the concept or utility of the invention. tions, authors, etc., if known.
For Sequence Searches Only Please incl appropriate serial number.	ude all pertinent information (parent, o	child, divisional, or issued patent numbers) along with the
Plene	Search S	Seg ID Nos
8,	10,25,42	, 43 x45
	please limit ?	
	please limit	
For Sig	4), plese	limit to size of GNTS
En Sy	{3 please)	limit suzi lo 6 NTS.
For Seg	45, please	e limit size to 6 NTS.
· ·	lese Search Lite In	Staterens + regular
	*********	*********
STAFF USE ONLY	Type of Search	Vendors and cost where applicable
Searcher:	214 0	STNDialog
Searcher Phone #:		Questel/Orbit Lexis/Nexis
Searcher Location:		Westlaw WWW/internet
		In-house sequence systems
Date Searcher Picked Up:	Gronographic	CommercialOligomerScore/Length
Date Completed:	Litigation	InterferenceSPDIEncode/Transl
Searcher Prep & Review Time:	Fulltext	Other (specity)
Online Time:	, Other	

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GenCore version 5.1.6
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July 20, 2005, 23:06:00; Search time 1253.4 Seconds (without alignments) 23.814 Million cell updates/sec
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/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq6:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq6:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq6:*
/ cgn2_6/ptodata/1/pna/USO8_NEW_COMB.seq2:*
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Maximum Match 100%
Listing first 100 summaries
OM nucleic - nucleic search, using sw model
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: /cgn2_6/ptodata/1/pna
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Gapop 10.0 , Gapext 1.0
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Perfect score:
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                                                                Run on:
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Query Match Length DB ID

Score

Result No.

Thu Jul 21 10:31:18 2005

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OM nucleic - nucleic search, using sw model

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Run on:

July 20, 2005, 22:36:35 ; Search time 1796 Seconds (without alignments) 136.380 Million cell updates/sec

US-09-735-363A-42 6 1 ggaagg 6 Title: Perfect score: Sequence:

Scoring table: IDENTITY_NUC Gapext 1.0

45554873 segs, 20411521753 residues

Searched:

13936 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 6

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 100 summaries

Pending Patents NA Main:* Database:

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July 20, 2005, 23:06:00; Search time 1253.4 Seconds (without alignments) 23.814 Million cell updates/sec
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/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq4:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq4:*
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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Maximum Match 100%
Listing first 100 summaries
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Gapop 10.0 , Gapext 1.0
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Query Match Length DB Score Result No.

OM nucleic - nucleic search, using sw model

Run on:

July 20, 2005, 22:36:35 ; Search time 1796 Seconds (without alignments) 136.380 Million cell updates/sec

US-09-735-363A-43 6 1 ggccgg 6

Title: Perfect score: Sequence:

IDENTITY NUC Gapop 10.0 , Gapext 1.0 Scoring table:

45554873 seqs, 20411521753 residues Searched:

13936

Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 6

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Pending_Patents_NA_Main:* Database :

Thu Jul 21 10:31:22 2005

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OM nucleic - nucleic search, using sw model

Run on:

July 20, 2005, 22:36:35 ; Search time 1796 Seconds (without alignments) 136.380 Million cell updates/sec

US-09-735-363A-45 6 1 999agg 6 Title: Perfect score: Sequence:

Scoring table:

IDENTITY NUC Gapop 10.0 , Gapext 1.0

45554873 seqs, 20411521753 residues Searched:

13936 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 6

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Database :

Pending Patents NA Main:*

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July 20, 2005, 23:06:00; Search time 1253.4 Seconds (without alignments) 23.814 Million cell updates/sec
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/cgn2_6/ptodate/1/pna/US11_NEW_COMB.seq3:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq4:*
/cgn2_6/ptodata/1/pna/US6_NEW_COMB.seq4:*
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0 NEW_COMB.seq4:
0 NEW_COMB.seq5:
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Maximum Match 100%
Listing first 100 summaries
OM nucleic - nucleic search, using sw model
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.: /cgn2_6/ptodata71/pna/PC
:: /cgn2_6/ptodata/1/pna/PC
:: /cgn2_6/ptodata/1/pna/US
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Gapop 10.0 , Gapext 1.0
                                                                                                                                                 US-09-735-363A-43
                                                                                                                                                                                                   1 ggccgg 6
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Perfect score:
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Description

SUMMARIES

Query Score Match Length DB

Result No. S

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GenCore version 5.1.6
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OM nucleic - nucleic search, using sw model
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July 20, 2005, 23:06:00; Search time 1253.4 Seconds (without alignments) 23.814 Million cell updates/sec US-09-735-363A-45 6. 1 gggagg 6 Title: Perfect score: Sequence:

Run on:

Scoring table:

IDENTITY NUC Gapop 10.0 , Gapext 1.0

16353451 segs, 2487343176 residues Searched:

2336 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 6

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Prodata1/pna/009. NEW_COMB.seq:*
/ptodata1/pna/US08 NEW_COMB.seq:*
/ptodata1/pna/US09 NEW_COMB.seq:*
/ptodata1/pna/US10 NEW_COMB.seq:*
/ptodata1/pna/US10 NEW_COMB.seq2:*
/ptodata1/pna/US10 NEW_COMB.seq2:*
/ptodata1/pna/US10 NEW_COMB.seq3:*
/ptodata1/pna/US10 NEW_COMB.seq4:*
/ptodata1/pna/US10 NEW_COMB.seq4:*
/ptodata1/pna/US10 NEW_COMB.seq6:* /cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq3:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq3:*
/cgn2_6/ptodata/1/pna/US11_NEW_COMB.seq4:*
/cgn2_6/ptodata/1/pna/US60_NEW_COMB.seq4:* /pna/PCT NEW COMB.seq: ptodata/1/pna/PCT Pending Patents NA N 1: /cgn2_6/ptodata/1 2: /cgn2_6/ptodata/1 3: /cgn2_6/ptodata/1 Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result.

Query Match Length DB Score

OM nucleic - nucleic search, using sw model

Run on:

July 20, 2005, 22:36:35 ; Search time 1796 Seconds (without alignments) 136.380 Million cell updates/sec

US-09-735-363A-10 6 1 gtgtgt 6 Title: Perfect score: Sequence:

Scoring table: IDENTITY_NUC Gapop 10.0, Gapext 1.0

45554873 segs, 20411521753 residues Searched:

13936 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 6

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Pending Patents NA Main:* Database :

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July 20, 2005, 23:06:00; Search time 1253.4 Seconds (without alignments) 23.814 Million cell updates/sec
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c/ptodata/l/pna/USOB_NEW_COMB.seq:*
c/ptodata/l/pna/USOB_NEW_COMB.seq:*
c/ptodata/l/pna/USIO_NEW_COMB.seq:*
c/ptodata/l/pna/USIO_NEW_COMB.seq2:*
c/ptodata/l/pna/USIO_NEW_COMB.seq2:*
c/ptodata/l/pna/USIO_NEW_COMB.seq3:*
c/ptodata/l/pna/USIO_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq6:*
c/ptodata/l/pna/USII_NEW_COMB.seq9:*
c/ptodata/l/pna/USII_NEW_COMB.seq3:*
c/ptodata/l/pna
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/cgn2_6/ptodata/1/pna/US60_NBW_COMB.seq:*
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GenCore version 5.1.6
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.: /cgn2_6/ptodata/1/pna/PCT
:: /cgn2_6/ptodata/1/pna/PCT
:: /cgn2_6/ptodata/1/pna/USC
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Maximum Match 100%
Listing first 100 summaries
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6
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence:
                                                                                                                                                                                                                                                                                                                                              Run on:
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Description

SUMMARIES

Query Match Length DB

Score

Result No.

OM nucleic - nucleic search, using sw model

July 20, 2005, 20:42:19 ; Search time 2266 Seconds (without alignments) 54.046 Million cell updates/sec Run on:

US-09-735-363A-8 3 1 gtg 3

Title: Perfect score: Sequence:

Scoring table:

IDENTITY NUC Gapop 10.0 , Gapext 1.0

45554873 seqs, 20411521753 residues Searched: 2500 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 3

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Pending_Patents_NA_Main:* Databasė :

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OM nucleic - nucleic search, using sw model
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July 20, 2005, 20:46:47; Search time 644 Seconds (without alignments) 23.168 Million cell updates/sec Title: Perfect score:

Run on:

US-09-735-363A-8

1 gtg 3 Sequence:

IDENTITY NUC Gapop 10.0 , Gapext 1.0 Scoring table:

16351682 segs, 2486654067 residues Searched:

132 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 3

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Database :

NEW COMB. Beg4: /cgn2_6/ptcdata/2/pna/US11_NEW_COMB.seq3:-/cgn2_6/ptcdata/2/pna/US11_NEW_COMB.seq4:-/cgn2_6/ptcdata/2/pna/US60_NEW_COMB.seq:* ptodata/2/pna/US10 ptodata/2/pna/US11 ptodata/2/pna/US11 Pending Patents NA New:* : /cgn2_6/ptodata/2/pna

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Query Score Match Length DB ID Result

OM nucleic - nucleic search, using sw model

July 20, 2005, 22:36:35 ; Search time 1796 Seconds (without alignments) 136.380 Million cell updates/sec Run on:

US-09-735-363A-25 6 1 999t99 6

Title: Perfect score: Sequence:

IDENTITY NUC Gapop 10.0 , Gapext 1.0 Scoring table:

45554873 segs, 20411521753 residues

Searched:

Total number of hits satisfying chosen parameters:

13936

Minimum DB seq length: 0 Maximum DB seq length: 6

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Post-processing: Minimum Match 0% Maximum Match 100% Listing first 100 summaries

Pending_Patents_NA_Main:* Database :